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## What is claimed is:

1. A drug dosage form comprising a compound susceptible to moisture-induced degradation and at least one pharmaceutically acceptable excipient prepared under conditions of low compression.

2. The drug dosage form of claim 1 comprising a capsule.

3. The drug dosage form of claim 1 comprising a capsule formed of hydroxypropyl methylcellulose.

4. The drug dosage form of claim 1 wherein the compound is contained in solid form within a capsule.

5. The drug dosage form of claim 1 wherein the form is subjected to no compression in excess of about 10,000 psi/g.

6. The drug dosage form of claim 1 wherein the form is subjected to no compression in excess of about 5,000 psi/g.

7. The drug dosage form of claim 1 wherein the form is subjected to no compression in excess of about 2,000 psi/g.

8. The drug dosage form of claim 1 wherein the excipient is hydroxypropyl methylcellulose, carboxymethyl cellulose, microcrystalline cellulose, amorphous silicon dioxide, magnesium stearate, starch, sodium starch glycolate, or a combination thereof.

9. The drug dosage form of claim 1 wherein the excipient has a residual moisture content of

less than about 10% by weight.

- 10. The drug dosage form of claim 1 exhibiting improved stability to moisture-induced degradation of the compound as compared with a tabletted form of the compound.
- 11. The drug dosage form of claim 1 comprising a unit dosage form.
- 12. A drug dosage form for a compound susceptible to moisture-induced degradation comprising the compound admixed with a substantially non-volatile, pharmaceutically acceptable oil.
- 13. The drug dosage form of claim 12 wherein the oil is an animal or vegetable oil.
- 14. The drug dosage form of claim 12 wherein said (oil is olive), corn, peanut, nut, soy, rapeseed, cottonseed, vitamin E, fish, or fallow-derived oil.
- 5. The drug dosage form of claim 12 wherein the oil is a mineral oil or silicone oil.
- 16. The drug dosage form of claim 12 wherein the compound oil admixture is present within a capsule.
- 17. The drug dosage form of claim 12/wherein the compound oil admixture is present within a soft shell capsule.
- 18. The drug dosage form of claim 12 wherein the compound oil admixture is present within a specially sealed hard-shell capsule.
- 19. The drug dosage form of claim 12 wherein at least some of the compound oil admixture

- is adsorbed on a pharmaceutically acceptable exciplent.
- 20. The drug dosage form of claim 12 wherein the excipient having the compound oil admixture adsorbed thereupon is within a capsule.
- 21. The drug dosage form of claim 12 wherein the excipient having the compound oil admixture adsorbed thereupon is within a table.
- 22. A drug dosage form for a compound susceptible to moisture-induced degradation comprising the drug and a pharmaceutically acceptable excipient admixed with a substantially non-volatile, pharmaceutically acceptable oil.
- 23. The drug dosage form of claim 22 wherein the oil is an animal or vegetable oil.
- 24. The drug dosage form of claim 22 wherein said oil is olive, corn, peanut, nut, soy, rapeseed, cottonseed, vitamin E, fish, or tallow-derived oil.
- 25. The drug dosage form of claim 22 wherein the oil is a mineral oil or silicone oil.
- 26. The drug dosage form of claim 22 wherein the drug and the excipient oil admixture are present within a capsule.
- 27. The drug dosage form of claim 22/wherein the drug and the excipient oil admixture are present within a tablet.
- 28. A drug dosage form comprising a compound susceptible to moisture-induced degradation admixed with a first pharmaceutically acceptable oil together with a pharmaceutically acceptable excipient admixed with a second pharmaceutically acceptable oil.

- 29. The drug dosage form of claim 28 wherein the first and the second pharmaceutically acceptable oils are, independently, an animal or vegetable oil.
- 30. The drug dosage form of claim 28 wherein the first and the second pharmaceutically acceptable oils are, independently, olive, corn, peanut, nut, soy, rapeseed, cottonseed, vitamin E, fish, or tallow-derived oil.
- 31. The drug dosage form of claim 28 wherein the first and the second pharmaceutically acceptable oils are, independently, a mineral oil or silicone oil.
- 32. The drug dosage form of claim 28 wherein the compound oil admixture and the excipient oil admixture are present within a capsule.
- 33. The drug dosage form of claim 28 wherein the compound oil admixture and the excipient oil admixture are present within a tablet.
  - A drug dosage form comprising a compound susceptible to moisture-induced degradation and at least one pharmaceutically acceptable hydrophobic powder.
- 35. The drug dosage form of claim 34 wherein the hydrophobic powder is triturated directly with the compound.
- 36. The drug dosage form of claim 34 wherein the hydrophobic powder is magnesium stearate.
- 37. A method for administering a compound susceptible to moisture-induced degradation to a patient comprising providing a unit dose of the compound which has not been processed

employing high compression.

- 38. A method for administering a compound susceptible to moisture-induced degradation to a patient comprising providing a unit dose of the compound admixed with a substantially non-volatile, pharmaceutically acceptable oil.
- 39. The method of claim 38 wherein the compound oil admixture is present within a capsule.
- 40. The method of claim 38 wherein the compound oil admixture is adsorbed on a pharmaceutically acceptable excipient.
- 41. A method for administering a compound susceptible to moisture-induced degradation to a patient comprising providing a unit dose of the compound and a pharmaceutically acceptable excipient admixed with a substantially non-volatile, pharmaceutically acceptable oil.
- 42. A method for administering a compound susceptible to moisture-induced degradation to a patient comprising providing a unit dose of the compound and at least one pharmaceutically acceptable hydrophobic powder.